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(54) Fischer-Tropsch catalysts containing iron and cobalt

(57) A process for preparing a Fischer-Tropsch catalyst comprises subjecting a slurry comprising a particulate alumina carrier, water and an active component selected from the group consisting in cobalt (Co), iron (Fe) and mixtures thereof, to a sub-atmospheric pressure

sure environment. The alumina carrier is thereby impregnated with the active component. The impregnated carrier is dried under a sub-atmospheric pressure environment. The dried impregnated carrier is calcined, thereby to obtain the Fischer-Tropsch catalyst.

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Description

THIS INVENTION relates to catalysis. It relates in particular to a process for preparing a Fischer-Tropsch catalyst, and to a Fischer-Tropsch catalyst prepared by the process.

According to the invention, there is provided a process for preparing a Fischer-Tropsch catalyst, which process comprises

subjecting a slurry comprising a particulate alumina carrier, water and an active component selected from the group consisting in cobalt (Co), iron (Fe) and mixtures thereof, to a sub-atmospheric pressure environment, thereby to impregnate the alumina carrier with the active component;

drying the impregnated carrier under a sub-atmospheric pressure environment; and
catching the dried impregnated carrier, thereby to obtain the Fischer-Tropsch catalyst.

The sub-atmospheric pressure environment during the impregnation may be at a pressure less than 20 kPa(a), and preferably at a pressure less than 10 kPa(a). Likewise, the sub-atmospheric pressure environment during the drying may be at a pressure less than 20 kPa(a), and preferably at a pressure less than 10 kPa(a).

The drying temperature is limited by the lower limit of the decomposition temperature of the active component, which is typically a nitrate salt so that the drying temperature is typically 70°C-90°C.

The sub-atmospheric pressure environments can thus be obtained by placing the slurry in a suitable enclosed vessel, and drawing the required sub-atmospheric pressure or vacuum on the vessel.

While the impregnation and drying in the sub-atmospheric pressure or vacuum environments or conditions can be effected in two separate or distinct steps, they can, if desired, be effected in a single step, so that the impregnation is effected while the drying takes place.

The drying in the sub-atmospheric pressure environment may be continued until the moisture content of the impregnated carrier is below 20% by mass. Thereafter, the impregnated carrier may be dried further under non-sub-atmospheric pressure conditions to remove more water, particularly water of crystallization. The further drying may be effected by passing a drying medium, eg air, in co-current or counter-current fashion over the impregnated carrier. The drying temperature may then be between 100°C and 180°C. Thus, for example, the further drying may be effected by means of hot air used to fluidize and dry the particulate carrier, eg in a tubular reactor, in which case the air flow is co-current. Instead, however, the further drying may be effected in a counter-current air drier, which may be a catalyst spray drier.

The calcination of the dried impregnated carrier thus converts or decomposes the active component to its oxide form. Thus, for example, the active component can be used in the form of a salt, eg $\text{Co}(\text{NO}_3)_2$, with the salt then being decomposed to the oxide of the active component, eg CO_3O_4 . The calcining is thus effected in a calciner. For example, the calciner can be mounted to the lower end of a spray drier used for further drying of the carrier as hereinbefore described, with the dried carrier then falling directly into the calciner.

If desired, the calcined catalyst may be re-slurried with water together with at least one of the following: the active component, another active component, or a dopant as hereinafter described, with the resultant impregnated carrier then again being subjected to drying and calcination, as hereinbefore described.

The process may include forming the slurry. In particular, the active component may initially be in the form of a water soluble compound of the component, and may then be dissolved in at least some of the water, prior to forming the slurry with the alumina carrier, so that formation of the slurry will then involve intimate admixing of the alumina carrier and a solution of the active component compound. Supersaturation during impregnation, which results in active component precursor crystallization, should be avoided during impregnation/drying. The supersaturation aspect is addressed through the slurry impregnation, while the vacuum drying at ~ 75°C of the aqueous solution addresses the precursor crystallization aspect. Thus, the purpose is to inhibit or prevent the diffusion of the catalyst precursor to the outer rim of the carrier body during drying (which would result in an egg-shell distribution) and which is enhanced by slow drying rates. Vacuum drying of an aqueous impregnation solution at ~ 75°C overcomes this problem, thereby also eliminating the option of using more volatile solvents, eg acetone, alcohol, etc, the use of which is also complicated by aspects such as: poorer solubilities of nitrates, for example ~ 35% less $\text{Co}(\text{NO}_3)_2$ is soluble in acetone as compared to water at room temperature; and the presence of high quantities of crystal waters, eg $\text{Co}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$.

While the alumina carrier will typically not be structurally promoted, it is, however, envisaged that it can contain a structural promoter such as magnesium (Mg) or cerium (Ce) if desired, eg if it is desired to enhance the attrition resistance of the resultant catalyst which is obtained from the process of the invention.

Irrespective of whether or not the alumina carrier is structurally promoted, the process of the invention may, however, be categorized thereby that no promoter to enhance the activity of the resultant catalyst or to modify its selectivity, such as potassium (K), chromium (Cr), magnesium (Mg), zirconium (Zr), ruthenium (Ru), thorium (Th), hafnium (Hf), cerium (Ce), rhenium (Re), uranium (U), vanadium (V), titanium (Ti), manganese (Mn), nickel (Ni), molybdenum (Mo),

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wolfram (W), lanthanum (La), palladium (Pd), uranium (U), praseodymium (Pr), neodymium (Nd) or other elements from groups IA or IIA of the periodic table of the elements, is added to the slurry or to the impregnated carrier. Thus, the resultant catalyst will then contain no such synthesis enhancing promoter(s). As a result, the calcination of the dried impregnated carrier may be effected at a relatively low temperature, eg at a temperature below 350°C, and even below 300°C.

When the catalyst is to be used in a slurry bed reactor, it may be washed with a suitable washing medium, eg water, after the calcination, to remove unwanted contaminants, such as cobalt, which may have formed on the external surface of the catalyst in the form of a shell of cobalt, ie without alumina being present in the shell. This washing is preferably effected with agitation, which may be achieved through boiling of the water in which the catalyst is washed. Changing the water from time to time speeds up the procedure.

The process may include reducing the calcined catalyst, eg by subjecting it to heat treatment under the influence of a reducing gas such as hydrogen.

It is usually desired that the resultant catalyst must comply geometrically with certain requirements in order to obtain a desired activity and/or selectivity, without the use of synthesis enhancing promoters, as hereinbefore described. Thus, for example, the catalyst may have a specified minimum pore size, typically a pore size of at least 12nm. If the alumina carrier geometry is such that those geometric requirements in respect of the resultant catalyst will not be met, then the process may include pretreating the alumina carrier appropriately. Thus, the process may include pretreating the particulate alumina carrier or substrate prior to forming the slurry thereof with the water and the active component, to modify the average diameter of its pores, ie its pore size, and/or to modify its chemical phase.

This pretreatment may comprise chemically pretreating the carrier and/or precalcining it prior to the slurry formation. When the carrier is chemically pretreated, this may involve treating it with ammonia. In particular, the ammonia treatment may comprise forming a paste by admixing the alumina carrier with water; spraying ammonia onto the paste; optionally, spraying more water onto the ammoniated paste, with simultaneous mixing, eg kneading, of the paste; extruding the paste; drying it; and then calcining it. This calcination may be effected at a temperature between 200°C and 1000°C, preferably between 500°C and 900°C. An acid, such as acetic acid, may be added to the paste, if desired.

When the carrier is precalcined without chemical pretreatment thereof as hereinbefore described, this calcination may also be effected at a temperature between 200°C and 1000°C, preferably between 500°C and 900°C. More particularly, the pretreatment may then comprise admixing the alumina carrier with water and an acid such as acetic acid; spraying additional water onto the mixture while mixing, eg kneading, it further; extruding the resultant paste; drying the extruded paste; and then effecting the calcination thereof. The water and acid initially mixed with the carrier may be in the form of dilute acid solution.

Naturally, the extrusion of the paste can be dispensed with if desired, eg if the resultant catalyst is to be used in a slurry bed reactor.

The alumina carrier or support may be that prepared by a spray-drying technique, provided that it has been subjected to the calcination temperature hereinbefore referred to, either during manufacture thereof, or subsequently during pretreatment thereof as hereinbefore described. Thus, a commercially available alumina support, such as the spray dried alumina support available from CONDEA Chemie GmbH of Überseering 40, 22297 Hamburg, Germany.

The alumina carrier is thus characterized thereby that it is used in a relatively pure form, containing at most only minor proportions of impurities or undesired substances such as titania and/or silica, and/or a minor proportion of a structural promoter as hereinbefore described. Furthermore, the process may be characterized thereby that the alumina carrier is the only carrier, ie that the alumina is not used in conjunction with other carriers or supports such as titania or silica.

The mass proportion of active component to alumina carrier in the slurry may be between 5:100 and 60:100, typically between 10:100 and 45:100.

The process may include adding to the slurry or to the impregnated uncalcined carrier or to the calcined catalyst, as a dopant, a minor proportion of an agent capable of enhancing the reducibility of the active component. The dopant may instead, or additionally, be added to the slurry which is formed when the calcined catalyst is reslurried as hereinbefore described. The dopant may comprise copper (Cu) and/or platinum (Pt). The mass proportion of the dopant, when present, to active component may be between 0.005:100 and 10:100, typically between 0.1:100 and 5.0:100 for copper, and between 0.01:100 and 0.3:100 for platinum.

The invention extends also to a Fischer-Tropsch catalyst, when produced by the process according to the invention. The catalyst has high specific activity, and is suitable for the selective conversion of synthesis gas, utilizing Fischer-Tropsch reaction conditions in fixed or slurry catalyst beds, to high molecular weight saturated hydrocarbons, ie waxes.

The invention will now be described in more detail with reference to the following non-limiting examples, and with reference to the accompanying drawings, in which

FIGURE 1 shows a plot of wax selectivity vs activity in respect of the catalysts of Examples 1 to 8;

FIGURE 2 shows a plot of wax selectivity vs pore size in respect of the catalysts of Examples 5, 7, 9, 10 and 11;

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FIGURE 3 shows a plot of wax selectivity vs pore size in respect of the catalysts of Examples 36 to 59; and FIGURE 4 shows a plot of activity vs selectivity in respect of the catalysts of Example 60.

In the examples hereunder, a series of cobalt supported catalysts on alumina were prepared and tested for their activity in the conversion of synthesis gas into hydrocarbons.

Fixed Bed Tests:

These tests were performed using 40mL of catalyst. The catalyst was either crushed and sieved extrudates to particle sizes ranging from 1mm to 1.7mm, or spray dried to particle sizes ranging between 0.05mm and 0.15mm. A tubular reactor was used, and had an internal diameter of 1cm and a length of 100cm. The top part of the reactor was filled with an inert material to act as a pre-heater for the gas feed. The feed gas consisted of hydrogen and carbon monoxide in an H₂/CO molar ratio of 2/1. The hydrogen and carbon monoxide accounted for about 84% (molar basis) of the feed. The other 16% was composed of inert gases, mainly methane (14.5%) and nitrogen (about 1%). The reactor was surrounded by an aluminium jacket which was electrically heated. The feed flow was controlled by means of Brooks mass flowmeters, and the Gas Hourly Space Velocity (GHSV) used in all the experiments was 4200 h⁻¹, based on total feed flow. The waxy products were collected in a condenser at about 18 bar and 130°C. This was followed by a condenser at about 18 bar and 20°C for the liquid products.

Slurry Phase Tests:

Between 10 and 30g of catalyst, spray-dried to particle sizes ranging between 38 µm to 150 µm, was suspended in 300mL molten wax and loaded in a CSTR with an internal volume of 500mL. The feed gas consisted of hydrogen and carbon monoxide in a H₂/CO molar ratio of 2/1. This reactor was electrically heated and sufficiently high stirrer speeds were employed so as to eliminate any gas-liquid mass transfer limitations. The feed flow was controlled by means of Brooks mass flow meters, and space velocities ranging between 1 and 3 m³/h/kg catalyst were used. GC analyses of the permanent gases as well as the volatile overhead hydrocarbons were used in order to characterize the product spectra.

All catalysts were reduced, prior to synthesis, in a fixed bed reactor at a pure hydrogen space velocity of 2 500 h⁻¹ and pressures ranging between 1 and 10 bar. The temperature was increased from room temperature to 950°C to 400°C at a rate of 1°C/min, after which isothermal conditions were maintained for 6 to 16 hours.

The catalysts were prepared according to the following examples:

Example 1

50g Alumina powder was added to 70mL distilled water. To this 50g Co(NO₃)₂·6H₂O was added. The mixture was kneaded thoroughly and extruded. The extrudates were dried in an oven for 2 to 3 hours at 100°C and thereafter calcined at 350°C for 16 hours. The alumina powder was that obtained from Degussa AG under the designation "Degussa Aluminium Oxide C".

Example 2

In a similar manner to Example 1, a catalyst was prepared by impregnation, drying and calcining, except that 42.5g, instead of 50g, Co(NO₃)₂·6H₂O was added to the alumina and water mixture.

Example 3

In a similar manner to Example 1, a catalyst was prepared but 37.5g, rather than 50g, Co(NO₃)₂·6H₂O was added to the alumina.

Example 4

In a similar manner to Example 1, a catalyst was prepared, but 20g Cr(NO₃)₃·9H₂O was added as a promoter.

Example 5

50g of the same alumina powder as used in Example 1, was added to 70mL distilled water. To this mixture 25g Co(NO₃)₂·6H₂O and 6.1g Mg(NO₃)₂·6H₂O were added. The mixture was kneaded and extruded similarly to Example 1.

$$\frac{2500 \text{ mL } H_2}{\text{hr mL cat}} \times \frac{1L}{\cancel{60 \text{ min}}} = \frac{41.67 \text{ mL } H_2}{\text{min mL cat}}$$

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Example 6

A catalyst was prepared in a similar manner to Example 1, but 0.35g KNO₃ was added as a promoter.

Example 7

A catalyst was prepared in a similar manner to Example 5, but 0.4g KNO₃ was added in place of the Mg (NO₃)₂·8H₂O.

Example 8

A catalyst was prepared in a similar manner to Example 1, but 4.9g Th(NO₃)₄·5H₂O was added as a promoter. The characteristics of the catalysts of Examples 1 to 8, as well as their performance in fixed bed Fischer-Tropsch synthesis, are presented in Table 1.

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TABLE 1

Catalyst Examples (Particle sizes varied between 1.0 and 1.7 mm with a pore size of 24 nm)	Active metal content (g Co per 100g Al ₂ O ₃)	Promoter Element	Fischer-Tropsch fixed bed synthesis performance at 18 bar and a GHSV of 4 200 h ⁻¹			
			Promotion level (expressed per 100g Al ₂ O ₃)	Reaction temperature (°C)	Fischer-Tropsch activity expressed as m mol H ₂ O formed per ml catalyst per h	
1	20	-	-	225	11.1	32
2	17	-	-	220	7.5	37
3	15	-	-	220	5.6	44
4	20	Cr	5.2 g	220	3.0	49
5	10	Mg	1.2 g	220	2.6	46
5	10	Mg	1.2 g	215	1.6	53
6	20	X	0.3 g	220	5.0	44
7	10	K	0.3 g	220	2.1	57
7	10	K	0.3 g	215	1.3	63
8	20	Ti	4.0 g	215	4.4	42

It can thus be seen that there is a strong correlation between the wax selectivity (defined here as the fraction of hydrocarbons condensed at 130°C at 18 bar) and the activity of the catalyst. This correlation is independent of the nature of the promoter and also independent of the addition of a promoter. This is more clearly indicated in Figure 1.

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which graphically shows the data of Table 1.

Additional supported cobalt catalysts were prepared according to the following procedure in order to cover a range of pore sizes.

5 Example 9

A catalyst was prepared in a similar manner to Example 1 but 12.5g $Mg(NO_3)_2 \cdot 6H_2O$ was added as a promoter.

10 Example 10

A catalyst was prepared in a similar manner to Example 5 except that 4.0g Zr(IV)acetylacetone was added in the place of the $Mg(NO_3)_2 \cdot 6H_2O$.

15 Example 11

A catalyst was prepared in a similar manner to Example 1, but 0.85g KNO_3 was added as a promoter. These catalysts were dried, calcined and tested for their fixed bed synthesis behaviour in a similar fashion to the catalysts of Examples 1 to 8. The physical characteristics and the catalytic activity of the catalysts are presented in table 2.

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TABLE 2

Catalyst Examples [Particle sizes varied between 1.0 and 1.7 mm]	Active metal content (g Co per 100g Al ₂ O ₃)	Promoter Element	Promotion level (expressed per 100g Al ₂ O ₃)	Platcar-Tropesch fixed bed synthesis performance at 18 bar, 220°C, and a GHSV of 4,200 h ⁻¹		Mass % reactor wax selectivity (~ C ₁₀)
				Pore size (nm)	Platcar-Tropesch activity expressed as in mol H ₂ O formed per mol catalyst per h	
9	20	Mg	2.48	21.5	2.5	24
10	10	Zr	1.58	22.5	2.1	30
5	10	Mg	1.28	24.0	2.4	46
7	10	X	0.38	24.3	2.1	57
11	20	K	0.78	25.9	2.3	61

From Table 2 it can thus be seen that for a given activity (ie ~ 2 mol H₂O/m² cat/h), reactor wax selectivity is a strong function of average catalyst pore size. This relationship is independent of the type of promoter added. This is more clearly illustrated in Figure 2, which graphically illustrates the results shown in Table 2.

In Examples 1 to 11, use was made of fumed Al₂O₃ which was co-extruded with the catalytically active components. An alternative approach is to extrude (or spray dry) and calcine the Al₂O₃ support separately as a first preparation step, prior to impregnation with the active component(s). This procedure allows for more freedom with respect to tailoring of the support geometry.

For this application, precipitated Al₂O₃, supplied by Condea Chemie GmbH, under their designations 'Pural SB alumina', 'Puralox SCCa 5/150, or Puralox HP 5/180' was used. The average pore size of the support was increased

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by the following pretreatment techniques: by calcination and/or by chemical treatment with an alkaline compound such as ammonia. Examples 12 to 35 thus are directed to pretreated supports.

5 Example 12

125mL acetic acid diluted with 1.7L distilled water was added to 2 kg Pural SB alumina powder obtained from Condea. Another 1.2L water was sprayed on while kneading the mixture. The alumina was extruded, dried at 120°C for 12 hours, and calcined at 600°C for 16 hours, to produce a pretreated support.

10 Example 13

A support was prepared in a similar manner to the support of Example 12, but using a calcination temperature of 700°C, instead of 600°C.

15 Example 14

A support was prepared in a similar manner to the support of Example 12, but using a calcination temperature of 800°C, instead of 600°C.

20 Example 15

125mL acetic acid diluted with 1.4L distilled water was added to 2 kg Pural SB alumina in a mixer. 250mL Ammonia (12.5 vol %) was sprayed onto this alumina paste. A further 1.2L water was sprayed onto the alumina while kneading the paste. The alumina was then extruded, dried at 120°C for 12 hours, and calcined at 600°C for 16 hours.

25 Example 16

A support was prepared in a similar manner to the support of Example 15, but using a calcination temperature of 700°C, instead of 600°C.

30 Example 17

A support was prepared in a similar manner to the support of Example 15, but using a calcination temperature of 800°C, instead of 600°C.

35 Example 18

19mL CH₃COOH was diluted to 210mL with distilled water. 20g of Zr(NO₃)₄·5H₂O was dissolved into this solution. This solution was then sprayed onto 300g Pural SB alumina while mixing in a mixer. 180mL of a 1.8 vol % ammonia solution was then sprayed onto the alumina while kneading the paste. The paste was then extruded, dried at 120°C for 2 hours and calcined at 750°C for 16 hours.

40 Example 19

A solution of 100g 4Mg(CO₃)₂·Mg(OH₂)₄·4H₂O, 160mL CH₃COOH and 150mL distilled water was sprayed onto 300g Pural SB alumina while mixing it in a mixer. This was then followed by spraying 220mL of 12.5 vol % ammonia onto the kneading mixture. After extruding the paste, the extrudates were dried at 120°C for 2 hours, and calcined at 750°C for 16 hours.

45 Example 20

A solution of 30g Zr(NO₃)₄·5H₂O in 210mL distilled water was sprayed onto 300g of Pural SB alumina while mixing in a mixer. While kneading this mixture, 180mL of a 3.5 vol % ammonia solution was sprayed onto it. The paste was then extruded, dried at 120°C for 2 hours, and calcined at 750°C for 16 hours.

50 Example 21

55 A support was prepared in a similar manner to the support of Example 18, but instead of 20g Zr(NO₃)₄·5H₂O, 30g

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Mg(NO₃)₂·6H₂O was used.

Example 22

5 A support was prepared in a similar manner to the support of Example 18, but instead of 20g Zr(NO₃)₄·5H₂O, 9g KNO₃ was used.

Example 23

10 A support was prepared in a similar manner to the support of Example 18, but instead of 20g Zr(NO₃)₄·5H₂O, 20g Mn(NO₃)₂·4H₂O was used.

Example 24

15 Puralox SCCa 5/150 support was calcined at 750°C for 16 hours.

Example 25

20 Puralox SCCa 5/150 support was calcined at 800°C for 16 hours.

Example 26

Puralox SCCa 5/150 support was calcined at 900°C for 16 hours.

25 Example 27

Puralox SCCa 5/150 support was calcined at 1 000°C for 16 hours.

Example 28

30 Puralox HP 5/180 support was calcined at 600°C for 16 hours.

Example 29

35 Puralox HP 5/180 support was calcined at 700°C for 16 hours.

Example 30

40 Puralox HP 5/180 support was calcined at 750°C for 16 hours.

Example 31

Puralox HP 5/180 support was calcined at 800°C for 16 hours.

45 Example 32

Puralox HP 5/180 support was calcined at 900°C for 16 hours.

Example 33

50 Puralox HP 5/180 support was calcined at 1 000°C for 16 hours.

Example 34

55 Puralox HP 5/180 support was calcined at 1 100°C for 16 hours.

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Example 35

A support was prepared in a similar manner to the support of Example 15, but using a calcination temperature of 750°C, instead of 800°C.

The physical properties of the pretreated supports of Examples 12 to 35 are given in Table 3.

TABLE 3

Example	Calcination Temperature °C	BET area m ² /g	Pore Volume mL/g	Average pore size nm
12	600	213	0.46	8.6
13	700	193	0.48	9.5
14	800	165	0.44	10.6
15	600	211	0.54	10.2
16	700	192	0.54	11.2
17	800	161	0.52	12.9
18	750	201	0.48	9.6
19	750	157	0.46	11.2
20	750	143	0.44	12.3
21	750	185	0.51	10.9
22	750	189	0.50	10.5
23	750	198	0.49	9.9
24	750	155	0.48	12.5
25	800	143	0.50	12.9
26	900	134	0.48	15.0
27	1 000	100	0.35	16.1
28	600	180	0.65	14.4
29	700	169	0.65	15.5
30	750	172	0.65	12.5
31	800	133	0.64	19.2
32	900	116	0.61	21.1
33	1 000	92	0.52	21.9
34	1 100	60	0.25	16.0
35	750	130	0.65	16.0

Increasing calcination temperature thus decreased the surface area of the supports. This effect was very similar for both types of support, i.e. with and without ammonia treatment.

The average pore size increased with an increase in the calcination temperature. The catalysts prepared with ammonia show a higher average pore size than the catalysts prepared in the absence of ammonia.

The supports of Examples 12 to 35 were impregnated with cobalt to determine the effect of their average pore size on wax selectivity. The following procedure was used:

50g of support was added to a solution of 50g Co(NO₃)₂·6H₂O and 0.05g Pt(NH₃)₄(NO₃)₂ in 50 to 70mL distilled water. The water was evaporated at 70°C under vacuum in a rotary evaporator. The catalyst was calcined at 250°C in a counter-current airflow for 6 hours.

The average pore sizes as well as reactor wax selectivities, as obtained in the tubular fixed bed reactor used in Examples 1 to 11, are shown in Table 4.

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TABLE 4

Sample	Example element	Promoter element	Average pore size (nm)	Mass & liquid hydrocarbons, drained as wax and oil (i.e. - C ₅ +) as measured during fixed bed reactor tests	
				Reaction conditions: Temperature : 200-208°C Pressure : 18 bar OHSV : 2600-3500 h ⁻¹	Vol% CO conversion: 10-20
36	12		7.5	64	62
37	13		8.0	52	52
38	14		8.9	57	57
39	15		6.6	60	60
40	16		5.7	63	63
41	17		10.7		
42	18		9.0	46	46
43	19	Hg	1.49	63	63
44	20	K	8.69	54	54
45	21	K	2.19	53	53
46	22	K	0.99	44	44
47	23	Mn	1.29	40	40
48	24		1.59	75	75
49	25			74	74
50	26			80	80
51	27			68	68
52	28			79	79
53	29			91	91
54	30			79	79
55	31			76	76
56	32			77	77
57	33			81	81
58	34			58	58
59	35				

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From Table 4 it can be seen that for a given activity, the reactor wax selectivity is a function of average catalyst pore size, independent of the type of promoter used (ie Zr, Mg, Mn, or K). This is more clearly illustrated in Figure 3, which summarizes the results shown in Table 4.

These tubular fixed bed synthesis Examples thus show that the main variables affecting wax selectivity from a cobalt based Fischer-Tropsch catalyst are the average pore size diameter of the support or carrier and the intrinsic catalyst activity.

In Examples 60 to 65 hereunder, commercially available spraydried and calcined Al_2O_3 Puralox SCCa 5/150 was used. This material was calcined at a temperature between 600°C and 700°C during manufacture thereof. This Al_2O_3 support material had a pore size of 12.5 nm which, as seen in Figure 3, is optimal from a wax selectivity and catalyst activity point of view. All the physical properties of this support material are listed in Table 5.

TABLE 5

Physical property	Al_2O_3 , Puralox SCCa 5/150 (used as selected support for the preparation of slurry phase Fischer-Tropsch catalysts)
Pore size	12.5 nm
Surface area	150 m ² /g
Pore volume	0.5 mL/g
Particle size distribution	45 - 150 μm
Crystalline phase	Gamma

Six catalyst samples were prepared with this support.

Example 60

40g $\text{Co}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ was dissolved in 50mL distilled water, and 50g Al_2O_3 Puralox SCCa 5/150 was suspended in this solution. This slurry was treated for ~ 2.5 hours at 75°C and 2 to 5 kPa in a rotary evaporator to impregnate the alumina carrier and to dry the impregnated carrier. The dried impregnated carrier was dried further and calcined at 230°C for 2 hours in an air flow of 1.5L/min. The resultant calcined sample was re-slurried in a solution that was made up by having dissolved 35g $\text{Co}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ and 50 mg $\text{Pt}(\text{NH}_3)_4(\text{NO}_3)_2$ in 50mL of distilled water. This slurry was again vacuum treated for ~ 2.5 hours at 75°C and 2 to 5 kPa in a rotary evaporator, followed by calcination at 230°C for 2 hours in an air flow of 1.5L/min.

Example 61

40g $\text{Co}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ was dissolved in 50mL distilled water, and 50g Al_2O_3 Puralox SCCa 5/150 was suspended in this solution. This slurry was treated for ~ 2.5 hours at 75°C and 2 to 5 kPa in a rotary evaporator to impregnate the alumina carrier and to dry the impregnated carrier. The dried impregnated carrier was calcined at 380°C for 5 hours in an air flow of 1.5L/min. The calcined sample was re-slurried in a solution that was made up by having dissolved 35g $\text{Co}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ in 50mL distilled water. This slurry was again vacuum treated for ~ 2.5 hours at 75°C and 2 to 5 kPa in a rotary evaporator, followed by calcination at 380°C for 5 hours in an air flow of 1.5L/min. The calcined sample was re-slurried in a solution that was made up by having dissolved 0.8g Flu (III) acetylacetone in 50mL acetone. This slurry was again vacuum treated, ie dried, until free flowing at 75°C and 2 to 5 kPa in a rotary evaporator, followed by a final calcination step at 330°C for 4 hours in an air flow of 1.5L/min.

Example 62

40g $\text{Co}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ and 1.2g perrhenic acid (HReO_4) was dissolved in 50mL distilled water, and 50g Al_2O_3 Puralox SCCa 5/150 was suspended in this solution. This slurry was vacuum treated for ~ 2.5 hours at 75°C in a rotary evaporator to impregnate the alumina carrier and to dry the impregnated carrier, followed by calcination at 350°C for 5 hours in an air flow of 1.5L/min. This calcined sample was re-slurried in a solution that was made up by having dissolved 35g $\text{Co}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ and 0.2g perrhenic acid in 50mL distilled water. This slurry was again vacuum dried for ~ 2.5 hours at 75°C until free flowing in a rotary evaporator, followed by calcination at 350°C for 5 hours in an air flow of 1.5L/min.

Example 63

29.6g $\text{Co}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ and 30 mg $\text{Pt}(\text{NH}_3)_4(\text{NO}_3)_2$ was dissolved in 50mL distilled water, and 50g Puralox SCCa

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5 S/150 was suspended in this solution. The slurry was vacuum treated for ~ 2.5 hours at 75°C and 2 to 5 kPa in a rotary evaporator to impregnate the alumina carrier and to dry the impregnated carrier. The dried impregnated carrier was calcined at 230°C for 2 hours in an air flow of 1.5L/min. The calcined sample was re-slurried in a solution that was made up by having dissolved 19.8g Co(NO₃)₂·6H₂O and 20 mg Pt(NH₃)₄(NO₃)₂ in 50mL of distilled water. This slurry was again vacuum dried for ~ 2.5 hours at 75°C and 2 to 5 kPa until free flowing in a rotary evaporator. The dried impregnated sample was calcined at 230°C for 2 hours in an air flow of 1.5L/min.

Example 64

10 This Example was similar to Example 61 with the following differences:

15 1st impregnation: 30g Co(NO₃)₂·6H₂O was used instead of 40g Co(NO₃)₂·6H₂O

20 2nd impregnation: 20g Co(NO₃)₂·6H₂O was used instead of 35g Co(NO₃)₂·6H₂O

25 3rd impregnation: 0.55 g Ru (III) acetylacetone was used instead of 0.8g Ru (III) acetylacetone

Thus, Examples 60 to 64 were prepared by means of slurry impregnation, ie impregnation solution in excess of the total available alumina carrier pore volume.

Example 65

25 26 kg Al₂O₃ Puralox SCCa S/150 was incipient impregnated with a 12.5L aqueous solution containing 13.9 kg Co(NO₃)₂·6H₂O and 8.6g Pt(NH₃)₄(NO₃)₂. This impregnated sample was dried at 80°C for 10 hours in an air flow of 40L/min, followed by calcination at 240°C for 4 hours in an air flow of 250L/min. In incipient impregnation, the volume of the impregnation solution used, is the aqueous solution referred to above, is equal to the pore volume of the alumina carrier.

30 A second incipient impregnation step followed during which this sample was impregnated with 11.3L of an aqueous solution containing 12.1 kg Co(NO₃)₂·6H₂O and 8.6g Pt(NH₃)₄(NO₃)₂. Drying and calcination was performed similarly to the first step.

35 A third and final incipient impregnation step followed during which this sample was impregnated with 13.2L of an aqueous solution containing 14.2 kg Co(NO₃)₂·6H₂O and 8.6g Pt(NH₃)₄(NO₃)₂, followed by the same drying and calcination steps as described above.

40 The preparation method of Example 60 was successfully scaled up to pilot plant scale, more or less on the same scale as that of Example 65. Proper vacuum drying proved to be an important parameter in the case of the scaled up version of the slurry impregnation option. The final moisture content of this dried impregnated catalyst should be less than ~ 20 mass %. This permits calcination where the dried impregnated catalyst is first passed through a counter current air drier (residence time of ~ 1min) set at 180°C, falling directly into a tubular calciner unit set at 250°C. The air flow through the calciner was set at ~ 8 dm³/kg cat/min at a superficial velocity of ~ 5 cm/s. Proper calcination required calcination periods in excess of 3 hours, preferably ~ 6 hours.

45 Examples 60, 63 and 65 were "unpromoted". Small quantities of Pt were added to assist with catalyst reduction. These quantities could vary between 0.03g Pt and 0.08g Pt per 100g Al₂O₃, and could be co-impregnated throughout all the impregnation steps (eg Example 65) or concentrated in the final impregnation step (eg Example 60).

The slurry phase Fischer-Tropel activities of catalyst Examples 60 to 65 are listed in Table 6.

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TABLE 6

Catalyst Sample	Active metal content (g Co per 100g Al ₂ O ₃)	Promoter	Fischer-Tropsch Slurry phase synthesis performance at 220°C, 20 bar, and at a space velocity of 2.0 m ³ /h/kg catalyst, (Feed gas: 33.3 vols CO and 66.7 vol % H ₂)				
			Element promotion level (expressed per 100g Al ₂ O ₃)	After 100 hours on line	After 400 hours on line	Vol % CO conversion	Productivity kg HC/kg Cat/h
60	30	Pt	0.05g	87	0.349	84	0.136
61	30	Ru	0.41g	77	0.307	70	0.281
62	30	Re	3.0g	70	0.281	NA	NA
63	20	Pt	0.05g	73	0.291	63	0.250
64	20	Ru	0.28g	73	0.288	63	0.252
65	31	Pt	0.05g	77	0.310	NA	NA

The following conclusions are evident from Table 6:

- Ru or Re promotion, which can be expensive at the required levels, does not result in enhanced specific Fischer-Tropsch activities at a cobalt content of ~ 20 mass % (ie 30g Co/100g Al₂O₃).

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- Applying a reported cobalt based Fischer-Tropsch kinetic equation, such as:

$$r_{FT} = (k_{FT} P_{H_2} P_{CO}) / (1 + \beta \cdot P_{CO})^2.$$

shows that intrinsic activity is linearly proportional to the cobalt content of a m Co/0.05 Pt/100 Al₂O₃ catalyst (Al₂O₃ Puralox SCCa 5/150) up to a level of m = 30 (ie constant cobalt utilization). At higher cobalt loadings (ie m > 30) cobalt utilization is diminished.

- In the preparation of the m Co/0.05 Pt/100 Al₂O₃ catalyst, the method of slurry impregnation (eg Example 60) is preferred with respect to incipient wetness impregnation (eg Example 65). The former impregnation method resulting in a catalyst with an intrinsic Fischer-Tropsch activity level ~ 1.35 times higher than the latter.

A selectivity investigation on this preferred cobalt slurry phase catalyst (ie Example 60) was performed and modelled. Table 7 provides an example of best fitted Schulz-Flory modelled selectivities of this catalyst, at representative synthesis conditions.

TABLE 7

% CO conversion at 220°C and 20 bar with a feed composed of 67 vol % H ₂ and 33 vol % CO	Mass % selectivities of the catalyst sample 36 after 400 hours on line				
	Fuelgas C ₁ -C ₂	LPG C ₃ -C ₄	Gasoline C ₅ -C ₁₁	Diesel C ₁₂ -C ₁₈	Wax C ₁₉ +
94	28	18	34	14	6
84	13	12	32	21	22
68	8	9	26	21	36
54	6	7	24	21	42
44	6	7	22	20	45
37	5	6	22	20	47
32	5	6	21	20	48
28	5	6	21	19	49
23	5	6	20	19	50
18	4	6	20	19	51
15	4	6	20	19	51

A graphical illustration of table 7 is presented in Figure 4, which underlines the dependence between activity and selectivity, as also deduced from figure 1 for the fixed bed application.

With respect to wax quality, slurry impregnation method (eg as described in the preparation of Example 60) is superior to the incipient wetness impregnation option (eg as described in the preparation of Example 65).

The reaction wax produced by catalyst Example 65, contained suspended sub-micron Co₃O₄ particles, at cobalt concentration level of ~ 100 ppm, which could not be removed by means of filtration through a Whatmans 42 filter paper. This also impacted negatively on the wax colour, and an undesirable saybolt colour of ~16 (ie darkest indicator) was determined for the filtered reactor wax. The origin of these sub-micron Co₃O₄ contaminant, was traced back to the presence of a clearly defined shell containing Co and no Al (~ 1 µm thick as observed through a SEM investigation), uniformly encapsulating the spray-dried γ Al₂O₃ spheres.

A thorough washing of the calcined catalyst Example 65, successfully removed this unwanted cobalt enriched material, without exerting any influence on the specific Fischer-Tropsch activity. This is despite of the fact that up to ~ 8% of the original cobalt content could be washed out.

Details of water washing procedure:

Experience gained during the washing of ~ 5 kg of catalyst Example 65 (ie after the final calcination step and before reduction), has shown that at least 25L of water is required per kg of catalyst.

Procedures that must be adhered to during the washing are:

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The water must be agitated to a limited degree, and this can be achieved through boiling.

- Changing water from time to time speeds up the procedure, eventually becoming clear, thus the recommended 25£ per kg of catalyst.

The unwanted situation of wax contamination has been proved to be almost absent in the case of slurry phase impregnated catalysts (eg sample 60), viz: catalysts with more homogeneous cobalt distribution throughout the particles, encapsulated by a far less pronounced cobalt oxide shell, is produced.

A water washing step is, however, still to be recommended in order to ensure a high quality wax. Wax produced by a washed slurry impregnated 30 Co/0.05 PV/100 Al₂O₃ catalyst contained only 1 to 3 ppm cobalt resulting in a saybolt colour of 10, after filtration through a Whatman's 42 filter paper.

Thus, very active cobalt based (fixed bed and slurry phase) Fischer-Tropsch catalysts can be prepared in a relatively inexpensive and easy manner, eg no expensive wax selectivity promoters are required in accordance with the invention.

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Claims

1. A process for preparing a Fischer-Tropsch catalyst, characterized in that the process comprises subjecting a slurry comprising a particulate alumina carrier, water and an active component selected from the group consisting in cobalt (Co), Iron (Fe) and mixtures thereof, to a sub-atmospheric pressure environment, thereby to impregnate the alumina carrier with the active component; drying the impregnated carrier under a sub-atmospheric pressure environment; and calcining the dried impregnated carrier, thereby to obtain the Fischer-Tropsch catalyst.
2. A process according to Claim 1, characterized in that the sub-atmospheric pressure environment during the impregnation is at a pressure less than 20 kPa(a).
3. A process according to Claim 1 or Claim 2, characterized in that the sub-atmospheric pressure environment during the drying is at a pressure less than 20 kPa(a).
4. A process according to any one of Claims 1 to 3 inclusive, characterized in that the impregnation and drying in the sub-atmospheric pressure environments is effected in a single step, so that the impregnation is effected while the drying takes place.
5. A process according to any one of Claims 1 to 4 inclusive, characterized in that the drying in the sub-atmospheric pressure environment is continued until the moisture content of the impregnated carrier is below 20% by mass, whereafter the impregnated carrier is dried further under non-sub-atmospheric pressure conditions by passing a drying medium in co-current or counter-current fashion over the impregnated carrier at a drying temperature between 100°C and 180°C.
6. A process according to any one of Claims 1 to 5 inclusive, characterized in that it includes adding to the slurry or to the impregnated uncalcined carrier, as a dopant, a minor proportion of an agent capable of enhancing the reducibility of the active component.
7. A process according to Claim 6, characterized in that the dopant comprises copper (Cu) and/or platinum (Pt), and wherein the mass proportion of the dopant to active component is between 0.005:100 and 10:100.
8. A process according to any one of Claims 1 to 7 inclusive, characterized in that the calcined catalyst is re-slurried with water together with at least one of the following: the active component, another active component, or, as a dopant, a minor proportion of an agent capable of enhancing the reducibility of the active component, with the resultant impregnated carrier then again being subjected to drying and calcination.
9. A process according to any one of Claims 1 to 8 inclusive, characterized in that the calcination of the dried impregnated carrier is effected at a temperature below 350°C.
10. A process according to any one of Claims 1 to 9 inclusive, characterized in that it includes washing the catalyst

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with a washing medium, to remove unwanted surface contaminants therefrom.

5 11. A process according to any one of Claims 1 to 10 inclusive, characterized in that it includes forming the slurry by dissolving a water soluble compound of the active component in water, prior to forming the slurry with the alumina carrier, and forming of the slurry by intimately admixing the alumina carrier and the solution of the active component compound.

10 12. A process according to Claim 11, characterized in that it includes pretreating the particulate alumina carrier prior to forming the slurry therewith with the water and the active component, to modify the average diameter of its pores and/or to modify its chemical phase, by chemically pretreating the carrier and/or precalcining the carrier prior to the slurry formation.

15 13. A process according to any one of Claims 1 to 12 inclusive, characterized in that the mass proportion of active component to alumina carrier in the slurry is between 5:100 and 60:100.

14. A Fischer-Tropsch catalyst characterized in that it is produced by the process according to any one of Claims 1 to 13 inclusive.

20 15. A method of preparing a high molecular weight saturated hydrocarbon wax which method comprises the selective conversion of synthesis gas under Fischer-Tropsch reaction conditions in the presence of a catalyst claimed in Claim 14.

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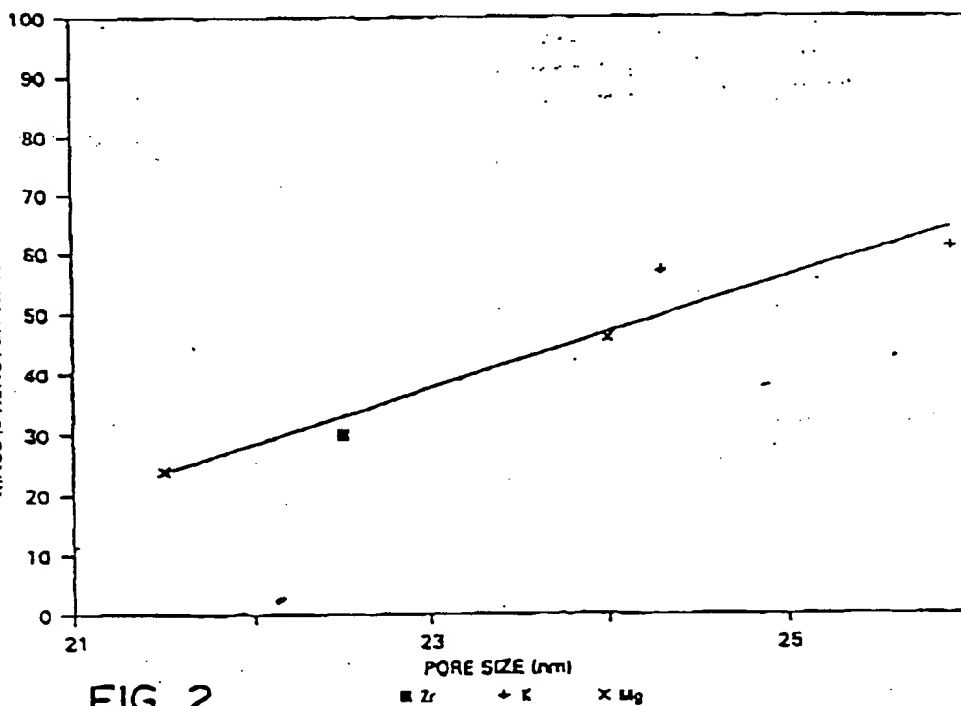
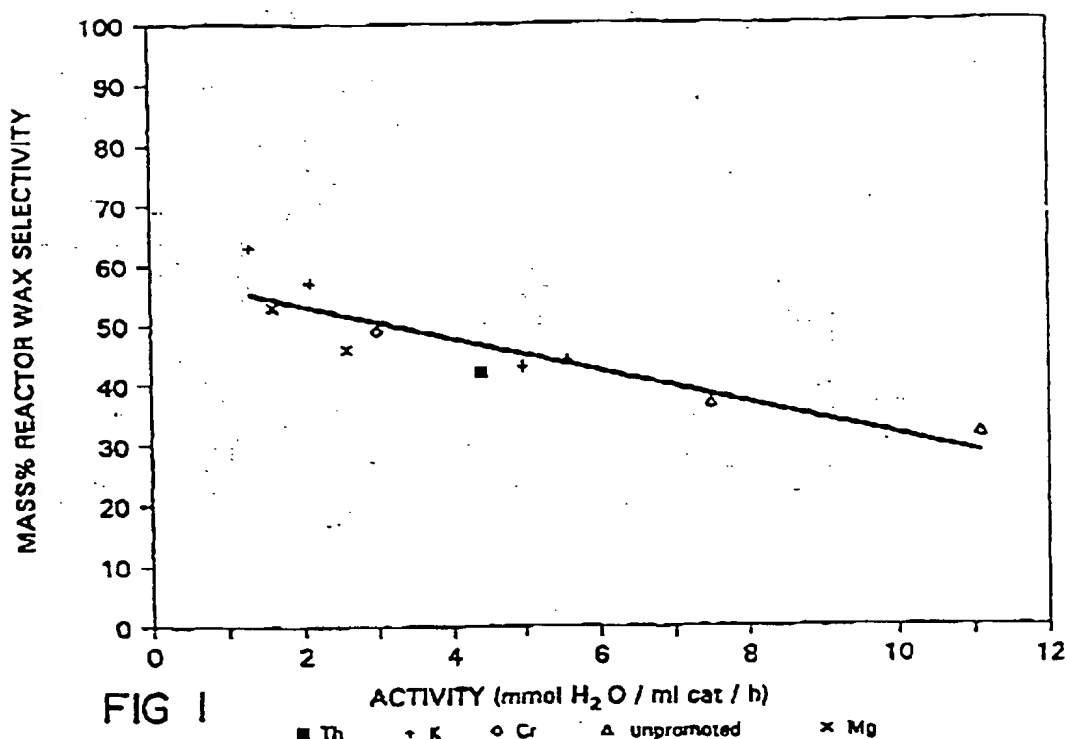


FIG 2

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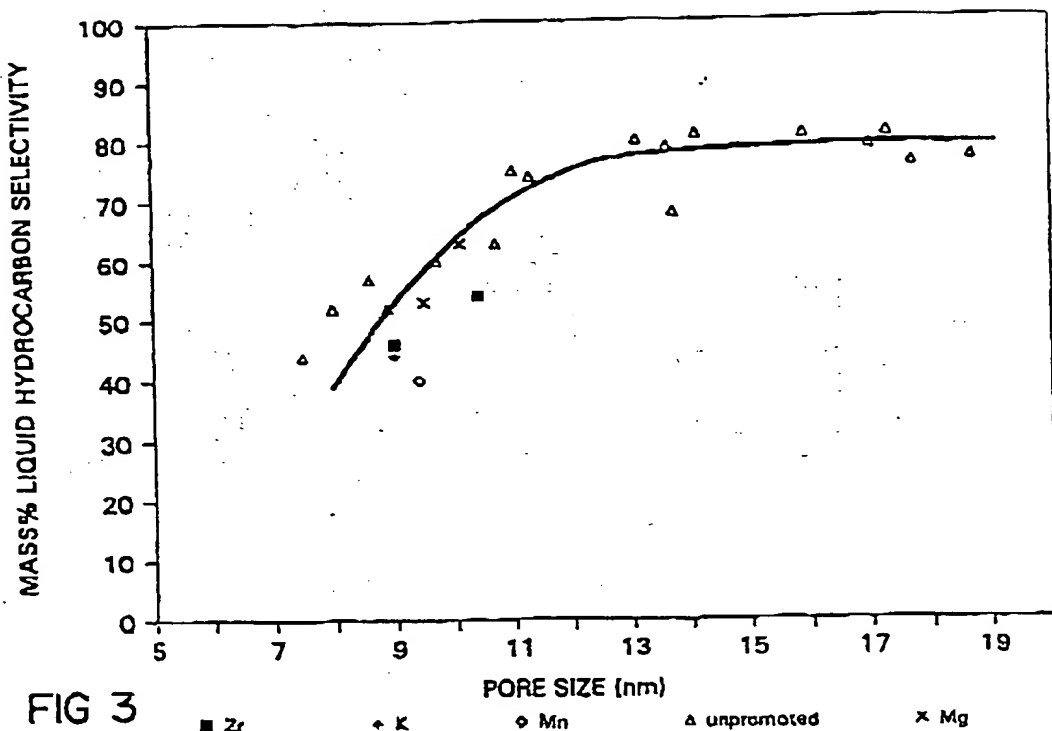


FIG 3

■ Zr + K ◊ Mn △ unpromoted x Mg

MASS% PRODUCT COMPOSITION

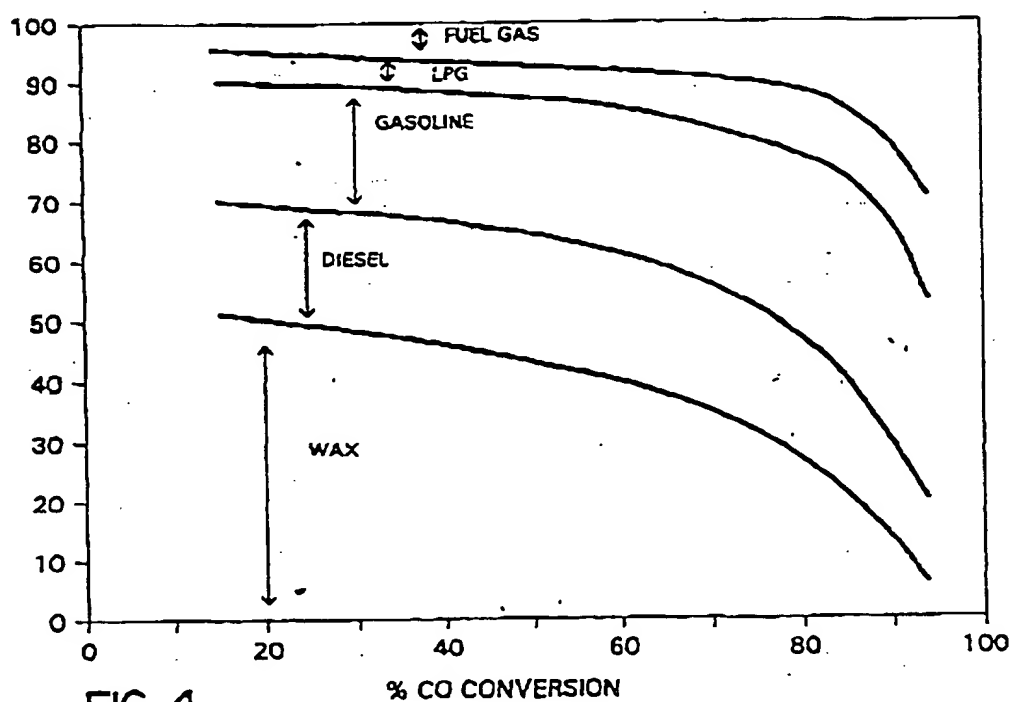


FIG 4

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European Patent
Office

EUROPEAN SEARCH REPORT

Application Number
EP 96 30 2437

DOCUMENTS CONSIDERED TO BE RELEVANT			CLASSIFICATION OF THE APPLICATION (EPO CLAS)
Category	Citation of document with indication, where appropriate, of relevant passages	Relevant to claim	
A	EP-A-8 535 798 (EXXON RESEARCH ENGINEERING CO) 7 April 1993 ---		B01J23/75 B01J23/745 C07C1/84
A	EP-A-0 434 284 (EXXON RESEARCH ENGINEERING CO) 26 June 1991 ---		
A	EP-A-0 266 898 (EXXON RESEARCH ENGINEERING CO) 11 May 1988 ---		
A	US-A-4 413 864 (BEUTHER HAROLD ET AL) 1 November 1983 ---		
A	EP-A-0 127 220 (SHELL INT RESEARCH) 5 December 1984 -----		
			TECHNICAL FIELDS SEARCHED (EPO CLAS)
			B01J C07C
The present search report has been drawn up for all claims			
Place of search	Date of completion of the search	Name of examiner	
THE HAGUE	18 July 1996	Thion, M	
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